



Diels Alder Reaction

Transcript

Instructor: Jessie Key

00:00:01:36 - 00:00:23:86

Instructor: Hello again, Doctor Jessie Key here. In this slide show, you'll be examining the Diels Alder reaction, a type of cycloaddition. The Diels Alder cycloaddition reaction is named after its discoverers, Otto Diels and Kurt Alder, two German chemists who were awarded the Nobel Prize in 1950, recognizing the importance of this reaction.

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Instructor: It is one of only a few carbon carbon bond forming reactions taught at the organic one and two level, and it is an important synthetic tool which has been employed in many syntheses, including the first synthesis of important biological molecules like prostaglandins. The Diels Alder reaction is an example of a four plus two cycloaddition, a type of paracyclic reaction. The four plus two cycloaddition classification is given as it involves two reagents, a four-carbon conjugated Pi system, the diene, and a two carbon Pi system, the dienophile, which come together to form a new substituted cyclohexene product.

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Instructor: Like all paracyclic reactions, the Diels Alder proceeds through a concerted single mechanistic step with a cyclic redistribution of bonding electrons. In this example, the Pi bond from the dienophile is going to form a new sigma bond between its uppermost vinyl carbon and the uppermost vinyl carbon of the diene. The uppermost pi bond of the diene comes down to form a new pi bond, and the lowermost pi bond of the diene goes to form a new sigma bond between the lowermost vinyl carbon of the diene and the lowermost vinyl carbon of the dienophile.

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Instructor: Note, when drawing the curved arrow notation mechanism for this reaction or any cycloaddition, the arrows can be drawn going around in a counterclockwise manner as shown or clockwise, the same results will be obtained. Examining a free energy diagram for this reaction with free energy on the y axis and reaction coordinate on the X axis, we can see this is a single reaction step with a cyclic transition state. The Diels-Alder reaction is in equilibrium with its reverse reaction, the retro Diels-Alder.

00:02:13:38 - 00:02:33:27

Instructor: Lower and moderate temperatures favor the forward reaction, while higher temperatures favor the retro Diels-Alder. The Diels Alder reaction is entropically unfavorable as

the reaction goes from two molecules to one molecule. Furthermore, that one molecule is cyclic which has less freedom.

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Instructor: However, breaking the three Pi bonds and formation of one new pi bond and two Sigma bonds is favorable in terms of enthalpy, since Sigma bonds are stronger than Pi bonds, resulting in a negative value for the change in enthalpy. We can therefore justify the reactivity observed for the reaction by taking into account the opposing enthalpic and entropic factors with the Gibbs free energy equation. ΔG is equal to ΔH plus negative $T \Delta S$. A lower value for

00:03:05:22 - 00:03:25:11

Instructor: the temperature will give a negative value for ΔG , allowing the DLS Alder reaction to proceed effectively. A higher value for temperature will give a positive value for ΔG , favoring the reverse reaction, the retro Diels-Alder. The structure of the dienophile plays a significant role in the usefulness of the reaction.

00:03:25:11 - 00:03:51:81

Instructor: Reactions using dienophiles with no electron withdrawing group present proceed very slowly due to having a relatively high activation energy. However, addition of an electron withdrawing group greatly reduces the activation energy and allows the reaction to proceed efficiently at lower temperatures. In this slide, the presence of the aldehyde group highlighted in red allows this Steel's alder reaction to proceed rapidly with approximately 100% yield.

00:03:51:81 - 00:04:12:98

Instructor: To help explain the effect of the electron withdrawing group, we can turn to a frontier molecular orbital theory explanation. Frontier molecular orbitals were first described by Japanese chemist Kenichi Fukui in 1952. For this important contribution to understanding organic reaction mechanisms, Fukui was later awarded a Nobel Prize in 1981.

00:04:12:98 - 00:04:38:11

Instructor: Frontier molecular orbital theory states that the most important molecular orbitals for chemical reactions are the frontier orbitals, the lUMO and HOMO. The lowest unoccupied molecular orbital abbreviated as lUMO is the orbital most capable of accepting electrons. The highest occupied molecular orbital abbreviated as HOMO is the orbital with electrons most able to react.

00:04:38:11 - 00:05:04:29

Instructor: In the case of the Diels-Alder reaction, the HOMO of the diene donates electrons to the lUMO of the dienophile. For a dienophile with no electron withdrawing group, there's a significant energy gap between these two molecular orbitals. However, when an electron withdrawing group is present on the dienophile, the energy level of the lUMO drops, decreasing the energy gap between the HOMO and lUMO, allowing the reaction to proceed more easily.

00:05:04:29 - 00:05:27:96

Instructor: Common electron withdrawing groups include carboxylic acids, esters, carbonyls, both aldehydes and ketones and nitriles. Carbonyl based electron withdrawing groups can withdraw electron density by both induction due to their electronegative oxygens and also by resonance. The dienophile can also dictate the stereochemistry of the product obtained.

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Instructor: When the dienophile is a *z* disubstituted alkene, the resulting cyclohexene product will have the two substituents arranged *cis*. However, when the dienophile is an *E* disubstituted alkene, a *trans* di substituted cyclohexene product will form. The conjugated diene structure also affects the Diels-Alder reactivity.

00:05:52:99 - 00:06:12:75

Instructor: Conjugate dienes can adopt one of two conformations, *S trans* and *SCIs*. This nomenclature refers to the arrangement of the two alkenes relative to the single bond connecting them. It is observed that only alkenes capable of adopting the *S* *cis* conformation undergo a Diels-Alder reaction.

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Instructor: This is because only the *S cis* conformation diene is capable of adopting the cyclic transition state with the dienophile, whereas the *trans* conformation carbons are too far apart in space to do this. This means that molecules unable to adopt the *S cis* conformation, like the three methyladine cyclohexene shown above, cannot participate in a Diels Alder reaction. Some molecules like cyclopentadiene are always in the *Sys* conformation and are more readily able to undergo the Diels Alder reaction.

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Instructor: In fact, cyclopentadiene is so reactive towards the Diels Alder reaction that even at room temperature, a molecule cyclopentadiene can react with another molecule of itself to form dicyclopentadiene. When cyclopentadiene and some other conjugated dienes are used, they can form bridged bicyclic products, having two possible stereochemical orientations of their substituents. *Endo* and *exo*.

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Instructor: The terms *exo* and *endo* refer to the relative position of the substituents relative to the bridge making the bicyclic system. Substituents which are *exo* are those which are *anti* or opposite to the larger of the bicyclic bridges. Substituents which are *endo* are those which are *syn* or on the same side as the larger bridge.

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Instructor: In this example, there is a larger two carbon bridge below a smaller one carbon bridge above. So, the *endo* substituent would be oriented relatively below, and the *exo* substituent would be oriented relatively above. Generally, the *endo* *pduct* is favored, often so much that it is the only product observed.

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Instructor: This is because in the cyclic transition state, there can be a favorable interaction which occurs between the electron withdrawing groups and the newly developing bonds. Looking at the energy diagram for this reaction with energy shown on the *y* axis and reaction progress on the *X* axis, the favorable interaction of the *endo* orientation lowers the energy of the transition state and therefore the activation energy. It is because of this lowered activation energy that the *endo* product tends to predominate, despite the greater stability seen for the *exo* substituted final product.